

GenCore version 5.1.6
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OM protein - protein search, using sw model

Run on: January 28, 2005, 13:12:07 ; Search time 154 Seconds
(without alignments)
270.212 Million cell updates/sec

Title: US-10-659-782A-32

Perfect score: 616

Sequence: 1 MFSPGTVCSLLGLMLWLDL.....PPSSRRSRHHQSPSCSPEL 116

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2002273 seqs, 358729299 residues

Total number of hits satisfying chosen parameters: 2002273

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

Database : A Geneseq_23Sep04:*

1: Geneseqp1980s:*

2: Geneseqp1990s:*

3: Geneseqp2000s:*

4: Geneseqp2001s:*

5: Geneseqp2002s:*

6: Geneseqp2003as:*

7: Geneseqp2003bs:*

8: Geneseqp2004s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	198	32.1	60	8	Adk66754 Human ghr
2	198	32.1	91	6	Aae33410 Human exo
3	198	32.1	117	2	Aaw87991 Protein d
4	198	32.1	117	3	Aay87236 Human sig
5	198	32.1	117	4	Aab20101 Z81933 pr
6	198	32.1	117	4	Aab26249 Human zsi
7	198	32.1	117	4	Aam38890 Human pol
8	198	32.1	117	4	Aab60511 Human ghr
9	198	32.1	117	5	Abb78319 Amino aci
10	198	32.1	117	5	Aae33838 Human zsi
11	198	32.1	117	5	Aae15883 Human zsi
12	198	32.1	117	6	Abu58046 Human PRO
13	198	32.1	117	6	Abu59124 Novel hum
14	198	32.1	117	6	Abu82636 Human sec
15	198	32.1	117	6	Abu17836 Novel hum
16	198	32.1	117	6	Abu60555 Human sec
17	198	32.1	117	6	Abu13937 Human PRO
18	198	32.1	117	6	Abu81090 Human PRO
19	198	32.1	117	6	Abu72522 Novel hum
20	198	32.1	117	6	Abu66790 Human PRO
21	198	32.1	117	6	Abu59871 Novel sec
22	198	32.1	117	6	Abu59271 Human sec
23	198	32.1	117	6	Abu25968 Human PRO
24	198	32.1	117	6	Abu25061 Human sec
25	198	32.1	117	6	Abu58977 Human sec

26	198	32.1	117	6	ABU92355	Abu92355 Novel hum
27	198	32.1	117	6	AAE33409	Aae33409 Human pre
28	198	32.1	117	6	ABU59420	Abu59420 Novel hum
29	198	32.1	117	6	ABU67066	Abu67066 Human sec
30	198	32.1	117	6	ABU92186	Abu92186 Novel hum
31	198	32.1	117	6	ABU10892	Abu10892 Human PRO
32	198	32.1	117	6	ABU81644	Abu81644 Novel hum
33	198	32.1	117	6	ABU88583	Abu88583 Human sec
34	198	32.1	117	6	ABO34097	Abu34097 Human PRO
35	198	32.1	117	6	ADA45961	Ada45961 Novel hum
36	198	32.1	117	6	ADA76392	Ada76392 Human PRO
37	198	32.1	117	6	ADA19042	Ada19042 Human PRO
38	198	32.1	117	6	ADA61665	Ada61665 Homo sapi
39	198	32.1	117	6	ADB19450	Adb19450 Novel hum
40	198	32.1	117	6	ADB27991	Adb27991 Human PRO
41	198	32.1	117	6	ADA86470	Ada86470 Novel hum
42	198	32.1	117	6	ADB16034	Adb16034 Human PRO
43	198	32.1	117	6	ADA37779	Ada37779 Human sec
44	198	32.1	117	6	ADA47820	Ada47820 Human PRO
45	198	32.1	117	6	ADA21465	Ada21465 Human sec

ALIGNMENTS

RESULT 1
ADK66754
ID ADK66754 standard; protein; 60 AA.
XX AC ADK66754;
XX
DT 06-MAY-2004 (first entry)
XX
DE Human ghrelin protein #1.
XX
XX Growth; appetite; fatness; genotype; polymorphism; ghrelin protein;
KW breeding; human.
XX
OS Homo sapiens.
XX
PN US2003211512-A1.
XX
PD 13-NOV-2003.
XX
PF 14-NOV-2002; 2002US-00294191.
XX
PR 14-NOV-2001; 2001US-0333222P.
XX
(ROTH/) ROTHSCCHILD M F.
(KIMK/) KIM K.
(ANDE/) ANDERSON L L.
XX
PI Rothechild MF, Kim K, Anderson LL;
XX
WPI; 2004-010657/01.
XX
Screening animals (i.e. pigs) to determine those more likely to produce
desired growth, appetite and fatness to optimize breeding and selection
techniques comprises detecting the presence of a polymorphism in the
Ghrelin gene.
XX
Disclosure; SEQ ID NO 3; 24pp; English.
XX
The present invention relates to a method of screening animals to
determine those more likely to produce desired growth, appetite and
fatness which involves obtaining a sample of genetic material from the
animal and assaying for the presence of a genotype in the animal which is
associated with favourable growth, appetite and fatness, the genotype
characterised by a polymorphism in the ghrelin gene. The composition and
methods are useful in screening animals (i.e. pigs) to determine those
more or less likely to produce desired growth, appetite and fatness to
optimise breeding and selection techniques. The present sequence is human
ghrelin protein of the invention.

```

XX SQ Sequence 60 AA;
Query Match 32.1%; Score 198; DB 8; Length 60;
Best Local Similarity 88.6%; Pred. No. 1.7e-14;
Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSTLLGLWLDLWDLAMAGSSFLSPHQVQVRPPHKAP 44
Db 1 MPSPGTVCSTLLGLWLDLWDLAMAGSSFLSPHQVQVRPPHKAP 44

RESULT 2
AAE33410
ID AAE33410 standard; protein; 91 AA.
XX
AC AAE33410;
XX
DT 02-APR-2003 (first entry)
DE Human exon 3-deleted ghrelin protein.
XX
KW Ghrelin; preproghrelin; GHS-R 1b; benign prostatic hyperplasia; therapy;
KW breast; cervical; uterine; choriocarcinoma; prostate; ovary; cytostatic;
KW cancer; human.
XX
OS Homo sapiens.
XX
PN WO200290387-A1.
XX
PD 14-NOV-2002.
XX
PF 10-MAY-2002; 2002WO-AU000582.
XX
PR 10-MAY-2001; 2001AU-00004919.
XX
PR 17-DEC-2001; 2001AU-00009567.
XX
PA (UYQU-) UNIV QUEENSLAND TECHNOLOGY.
XX
PI Chopin LK, Jeffery PL, Herington AC;
XX
WPI; 2003-111957/10.
DR N-PSDB; AAD50726.
XX
XX Identifying a cancer cell or tissue for treating prostate, ovarian,
PT breast cancer, or benign prostatic hyperplasia, by detecting the
PT expression of a ghrelin, an exon-3 deleted preproghrelin and/or a GHS-R
PT 1b proteins or nucleic acids.
XX
PS Claim 14; Page 34; 50pp; English.
XX
CC The invention relates to a method for identifying a cancer cell or tissue
CC of the reproductive system by detecting expression of a ghrelin, an exon-
CC 3 deleted preproghrelin and/or a GHS-R 1b proteins or nucleic acids. The
CC antibodies, exon 3-deleted form of preproghrelin and antagonists are
CC useful for treating cancer of the reproductive system such as prostate,
CC ovarian, breast, cervical or uterine cancer, choriocarcinoma or benign
CC prostatic hyperplasia. The present sequence is human exon 3-deleted
CC ghrelin protein
XX
SQ Sequence 91 AA;
Query Match 32.1%; Score 198; DB 6; Length 91;
Best Local Similarity 88.6%; Pred. No. 2.8e-14;
Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSTLLGLWLDLWDLAMAGSSFLSPHQVQVRPPHKAP 44
Db 1 MPSPGTVCSTLLGLWLDLWDLAMAGSSFLSPHQVQVRPPHKAP 44

RESULT 3
AAW87991
ID AAW87991 standard; protein; 117 AA.
XX
AC AAW87991;
XX
DT 07-APR-1999 (first entry)
DE Protein designated zsig33.
XX
KW Zsig33; gastric motility; gastrointestinal inflammation; reflux disease;
KW nutrient absorption regulation; obesity; metabolic disorder.
XX
OS Homo sapiens.
XX
Key Location/Qualifiers
FH Peptide 1..23
FT /note= "signal peptide"
FT Protein 24..117
FT /note= "mature protein"
XX
PN WO9842840-A1.
XX
PD 01-OCT-1998.
XX
PF 23-MAR-1998; 98WO-US0005620.
XX
PR 24-MAR-1997; 97US-0041102P.
PR 24-MAR-1997; 97US-00822897.
XX
PA (ZYMO ) ZYMOGENETICS INC.
XX
PI Sheppard PO, Deisher TA;
XX
WPI; 1999-070071/06.
DR N-PSDB; AAX04550.
XX
XX Human polypeptide having homology to motilin, zsig33 - useful e.g. to
PT treat gastrointestinal motility disorders, obesity etc. and to identify
PT antagonists to treat gastrointestinal hypermotility.
XX
PS Claim 13; Page 55-56; 69pp; English.
XX
CC The present sequence represents a protein designated Zsig33. The nucleic
CC acids are strongly expressed in stomach tissue. The polypeptide (or
CC allelic variants/orthologs) can be used to stimulate gastric motility,
CC measured as increased transit time or gastric emptying of an ingested
CC substance in mammals. The products are used to treat disorders associated
CC with gastrointestinal cell contractility, secretion of digestive
CC enzymes/acids, gastrointestinal motility, recruitment of digestive
CC enzymes, gastrointestinal inflammation, reflux disease and nutrient
CC absorption regulation. Zsig33 polypeptides may also be important
CC neurologically, since the family of gut-brain peptides to which the
CC homologous protein motilin belongs has been associated with neurological
CC and CNS functions. They may therefore be used e.g. to regulate satiety or
CC treat obesity and other metabolic disorders where neurological feedback
CC modulates nutritional absorption. They are useful to identify zsig33
CC agonists, antagonists and ligands and to produce antibodies
XX
SQ Sequence 117 AA;
Query Match 32.1%; Score 198; DB 2; Length 117;
Best Local Similarity 88.6%; Pred. No. 3.9e-14;
Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSTLLGLWLDLWDLAMAGSSFLSPHQVQVRPPHKAP 44
Db 1 MPSPGTVCSTLLGLWLDLWDLAMAGSSFLSPHQVQVRPPHKAP 44

RESULT 4
AAW87236
ID AAW87236 standard; protein; 117 AA.
XX
AC AAW87236;

```

XX DT 11-MAY-2000 (first entry)
XX DE Human signal peptide containing protein HSP-13 SEQ ID NO:13.
XX DE
XX DE
XX KW Human; signal peptide-containing protein; HSP; diagnosis; cancer;
KW inflammation; cardiovascular disease; anticancer; anti-inflammatory;
KW antimicrobial; neurotropic; neuroprotective; cardiovascular; hepatotropic;
KW antiasthmatic; gene therapy; cell proliferation; neurological disorder;
KW reproductive disorder; developmental disorder; arteriosclerosis;
KW cirrhosis; psoriasis; acquired immune deficiency syndrome; anaemia;
KW asthma; Crohn's disease; infection; Alzheimer's disease; schizophrenia;
KW Parkinson's disease; Huntington's diseases; ovulatory defect;
KW muscular dystrophy.
XX OS Homo sapiens.
XX PN WO200000610-A2.
XX PD 05-JAN-2000.
XX PF 25-JUN-1999; 99WO-US014484.
XX PR 26-JUN-1998; 98US-0090762P.
XX PR 31-JUL-1998; 98US-0094983P.
XX PR 01-OCT-1998; 98US-0102686P.
XX PR 11-DEC-1998; 98US-0112129P.
XX PA (INCY-) INCYTE PHARM INC.
XX PI Lal P, Tang YT, Gorgone GA, Corley NC, Guegler KJ, Baughn MR;
PI Akerblom IE, Au-Young J, Yue H, Patterson C, Reddy R, Hillman JL;
PI Bandman O;
XX WPI; 2000-160673/14.
XX N-PSDB; AA298121.
XX
XX New human signal peptide-containing proteins useful in treatment,
PT prevention and diagnosis of-e.g. cancer, inflammation and cardiovascular
PT disease.
XX
XX Claim 1; Page 168-169; 327pp; English.
XX
XX AA298109 to AA298242 encode AA297224 to AA297357 which represent the
CC human signal peptide-containing proteins HSP-1 to HSP-134. HSPs have
CC anticancer, anti-inflammatory, antimicrobial, neurotropic, hepatotropic,
CC neuroprotective, cardiovascular and antiasthmatic activities, and can be
CC used in gene therapy. HSPs can be used to treat or prevent disorders
CC associated with decreased activity or function of HSP. Antagonists of
CC HSP are used to treat or prevent disorders associated with increased
CC activity or function of HSP. Such diseases include cell proliferation
CC (including cancer), inflammation, cardiovascular, neurological,
CC reproductive or developmental disorders, (e.g. arteriosclerosis,
CC cirrhosis, psoriasis, acquired immune deficiency syndrome, anaemia,
CC asthma, Crohn's disease, microbial or other infections, congestive or
CC ischaemic heart disease, Alzheimer's, Parkinson's or Huntington's
CC diseases, schizophrenia, ovulatory defects, muscular dystrophy). HSP
CC nucleic acids can be used for the recombinant production of HSP, for
CC detecting HSP in standard hybridisation and amplification assays (for
CC diagnosis and monitoring), in gene therapy, as antisense, triplex-forming
CC or ribozyme therapeutics, for detecting related sequences or genetic
CC variations, and for chromosomal mapping. HSP are also used to raise
CC specific antibodies (Ab) and to screen for agonists and antagonists
CC (potential therapeutic agents). Ab are used to diagnose, or monitor, HSP
CC -related diseases (in usual immunoassays), as therapeutic antagonists, in
CC competitive drug screens, and for purification of HSP from natural
CC sources
XX SQ Sequence 117 AA;

Query Match 32.1%; Score 198; DB 3; Length 117;
Best Local Similarity 88.6%; Pred. No. 3.9e-14;
Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
DB 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
RESULT 5
AAB20101
ID AAB20101 standard; protein; 117 AA.
XX AC AAB20101;
XX DT 23-APR-2001 (first entry)
XX DE Zeig33 protein.
XX KW SGIP; zaig33; anorectic; antidiabetic; somatotropin; somatomedin-C;
KW nutritional absorption modulator; growth hormone secretagogue; therapy;
KW human.
XX OS Homo sapiens.
XX FH Key Location/Qualifiers
FT Peptide 1..23 /label= Signal_peptide
FT Protein 24..117 /label= Mature_protein
FT Peptide 24..34 /label= SGIP_peptide
FT /note= "this peptide is claimed in Claim 1"
XX WO200100830-A1.
XX PN 04-JAN-2001.
XX PD 30-JUN-2000; 2000WO-US018306.
XX PF 30-JUN-1999; 99US-00345157.
XX PR (ZYMO) ZYMOGENETICS INC.
XX PA Sheppard PO, Jaspers SR, Deisher TA, Bishop PD;
XX PI WPI; 2001-123010/13.
XX DR N-PSDB; AAF30033.
XX Novel variants of SGIP peptides for modulating contractility in duodenum
PT or jejunum tissue, pancreatic secretion of hormones and digestive
PT enzymes, inducing growth hormone secretion or modulating gastric
PT emptying.
XX Disclosure; 54; 61pp; English.
XX The present sequence is that of zaig33, a secreted protein with homology
CC to motilin (see AAB20102). Zeig33 is expressed at high levels in the
CC stomach, and at lower levels in the small intestine and pancreas. A novel
CC peptide fragment of zaig33, termed SGIP (see AAB20100), is claimed. SGIP
CC is a ligand for growth hormone secretagogue receptor, and is therefore
CC useful for modulating secretion of growth hormone and insulin like growth
CC factor 1. SGIP, and variant SGIP peptides, are used in claimed methods
CC for stimulating contractility in duodenum or jejunum tissue, modulating
CC pancreatic secretion of hormones and digestive enzymes, inducing growth
CC hormone secretion, and modulating gastric emptying
XX SQ Sequence 117 AA;

Query Match 32.1%; Score 198; DB 4; Length 117;
Best Local Similarity 88.6%; Pred. No. 3.9e-14;
Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
DB 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44

RESULT 6
 AAB62649
 ID AAB62649 standard; protein; 117 AA.
 XX
 AC AAB62649;
 DT 23-JUL-2001 (first entry)
 XX
 DE Human zsig33 polypeptide.
 XX
 KW zsig33; signal transduction; hormone; enzyme; neural development;
 KW gastric contractility; nutrient uptake; digestive; pancreatic; human;
 KW insulin-like growth factor-I; growth hormone; bone; gastrointestinal;
 KW glucose; osteopathic; anorectic; vulnerable; immunomodulator; GHS-R;
 KW G-protein coupled receptor.
 XX
 OS Homo sapiens.
 XX
 FH Key Location/Qualifiers
 FT Peptide 24..37
 FT /note= "specifically claimed fragment that binds to the
 FT GHS-R"
 FT
 XX
 WO200138355-A2.
 XX
 31-MAY-2001.
 XX
 22-NOV-2000; 2000WO-US032074.
 XX
 22-NOV-1999; 99US-0166765P.
 XX
 (ZYMO) ZYMOGENETICS INC.
 XX
 PI Sheppard PO, Jaspers SR, Deisher TA, Bishop PD;
 XX
 WPI; 2001-355879/37.
 DR N-PSDB; AAF83678.
 XX
 Forming reversible peptide receptor complex for purifying cell and
 PT peptides, stimulating signal transduction and modulating hormone
 PT secretion, involves contacting a receptor with zsig33 polypeptide.
 XX
 PS Claim 1; Page 93-94; 111pp; English.
 XX
 CC The invention relates to a method of forming a reversible peptide-
 CC receptor complex that involves providing an immobilized receptor, and
 CC contacting the receptor with a zsig33 peptide (comprising residues 24-37
 CC of AAB62649), where the receptor binds to the zsig33 peptide. The method
 CC is useful for purifying cells, purifying a peptide, stimulating signal
 CC transduction in a cell expressing a receptor. It is also useful for
 CC modulating secretion of hormones, neural development and/or utilization,
 CC gastric contractility, nutrient uptake, secretion of digestive and
 CC pancreatic enzymes and hormones, secretion of insulin-like growth factor
 CC -I, secretion of non-zsig33 proteins. It is useful for modulating growth
 CC hormone secretion in a mammal having a disease associated with abnormal
 CC levels of growth hormone, such as osteoporosis, bone repair, bone
 CC remodeling, low osteoblast levels, cartilage repair and remodeling,
 CC skeletal dysplasia, immune suppression, obesity, growth retardation,
 CC protein catabolic responses after surgery, cachexia, protein loss,
 CC dwarfism, wound healing and ovulation induction, treating a mammal having
 CC a metabolic disorder requiring neurological feedback, such as satiety
 CC regulation, glucose absorption and metabolism and neuropathy-associated
 CC gastrointestinal disorders, and stimulating glucose-induced insulin
 CC release in a mammal. The present sequence represents the human zsig33
 CC polypeptide, a peptide ligand for the G-protein coupled receptor, GHS-R
 XX
 SQ Sequence 117 AA;
 Query Match 32.1%; Score 198; DB 4; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;
 Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

OY 1 MPSPGTVCSSLLLLGMLWLDLAWAGSSFLSPHQRVQVRPPHKAP 44
 |||||
 DB 1 MPSPGTVCSSLLLLGMLWLDLAWAGSSFLSPHQRVQVRPPHKAP 44
 |||||
 RESULT 7
 AAM38890
 ID AAM38890 standard; protein; 117 AA.
 XX
 AC AAM38890;
 XX
 DT 22-OCT-2001 (first entry)
 XX
 DE Human polypeptide SEQ ID NO 2035.
 XX
 KW Human; nootropic; immunosuppressant; cytostatic; gene therapy; cancer;
 KW peripheral nervous system; neuropathy; central nervous system; CNS;
 KW Alzheimer's; Parkinson's disease; Huntington's disease; haemostatic;
 KW amyotrophic lateral sclerosis; Shy-Drager Syndrome; chemotactic;
 KW chemokinetic; thrombolytic; drug screening; arthritis; inflammation;
 KW leukaemia.
 XX
 OS Homo sapiens.
 XX
 PN WO200153312-A1.
 XX
 26-JUL-2001.
 XX
 26-DEC-2000; 2000WO-US034263.
 XX
 23-DEC-1999; 99US-00471275.
 PR 21-JAN-2000; 2000US-00488725.
 PR 25-APR-2000; 2000US-00552317.
 PR 20-JUN-2000; 2000US-00598042.
 PR 19-JUL-2000; 2000US-00620312.
 PR 03-AUG-2000; 2000US-00653450.
 PR 14-SEP-2000; 2000US-00662191.
 PR 19-OCT-2000; 2000US-00693036.
 PR 29-NOV-2000; 2000US-00727344.
 XX
 (HYSE-) HYSEQ INC.
 XX
 PI Tang YT, Liu C, Asundi V, Chen R, Ma Y, Qian XB, Ren F, Wang D;
 PI Wang J, Wang Z, Wehrman T, Xu C, Xue AJ, Yang Y, Zhang J, Zhao QA;
 PI Zhou P, Goodrich R, Drmanac RT;
 XX
 WPI; 2001-442253/47.
 DR N-PSDB; AAI58046.
 XX
 Novel nucleic acids and polypeptides, useful for treating disorders such
 PT as central nervous system injuries.
 XX
 Example 3; SEQ ID NO 2035; 10078pp; English.
 PS
 XX The invention relates to human nucleic acids (AAI57798-AAI61369) and the
 CC encoded polypeptides (AAM38842-AAM42213) with nootropic,
 CC immunosuppressant and cytostatic activity. The polynucleotides are useful
 CC in gene therapy. A composition containing a polypeptide or polynucleotide
 CC of the invention may be used to treat diseases of the peripheral nervous
 CC system, such as peripheral nervous injuries, peripheral neuropathy and
 CC localised neuropathies and central nervous system diseases, such as
 CC Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic
 CC lateral sclerosis, and Shy-Drager Syndrome. Other uses include the
 CC utilisation of the activities such as: Immune system suppression,
 CC Activin/inhibin activity, chemotactic/chemokinetic activity, haemostatic
 CC and thrombolytic activity, cancer diagnosis and therapy, drug screening,
 CC assays for receptor activity, arthritis and inflammation, leukaemias and
 CC C.N.S disorders. Note: The sequence data for this patent did not form
 CC part of the printed specification
 XX
 SQ Sequence 117 AA;

Query Match 32.1%; Score 198; DB 4; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;
 Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||
 DB 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||

RESULT 8
 AAB60511
 ID AAB60511.standard; protein; 117 AA.
 XX
 AC AAB60511;
 XX
 DT 24-APR-2001 (first entry)
 XX
 DE Human ghrelin preproprotein, SEQ ID NO:5.
 XX
 KW Growth hormone secretagogue; GHS; ghrelin; precursor; preproprotein;
 KW calcium concentration elevation; infant growth disorder;
 KW growth hormone deficiency.
 XX
 OS Homo sapiens.
 XX
 PN WO200107475-A1.
 XX
 PD 01-PEB-2001.
 XX
 PF 24-JUL-2000; 2000WO-JP004907.
 XX
 PR 23-JUL-1999; 99JP-00210002.
 PR 29-NOV-1999; 99JP-00338841.
 PR 26-APR-2000; 2000JP-00126623.
 XX
 PA (KANG/) KANGAWA K.
 XX
 PI Kangawa K, Kojima M, Hosoda H, Matsuo H, Minamitake Y;
 XX
 DR WPI: 2001-159704/16.
 DR N-PSDB; AAF59645.
 XX
 PS New peptide compounds which induce growth hormone secretion and elevate
 PT cell calcium concentrations, useful in treatment and diagnosis of infant
 PT growth disorders.
 XX
 PS Claim 3; Page 182; 210pp; Japanese.
 XX
 CC The invention relates to a novel peptide compound or its salt which
 CC induces the secretion of growth hormone and/or elevates calcium ion
 CC concentration in cells. The peptides are ghrelin homologues and are
 CC characterised in that at least one amino acid has been substituted by a
 CC modified amino acid and/or a non-amino acid compound. The invention also
 CC encompasses the unmodified peptides; the DNA encoding the peptides;
 CC vectors and host cells comprising such DNA; a method of producing the
 CC peptides comprising recombinant production, optionally followed by
 CC chemical modification; an antibody specific for a peptide of the
 CC invention; and an assay and kit for detecting the peptides. The peptides
 CC of the invention are useful for treating and/or diagnosing diseases
 CC caused by a deficiency in growth hormone expression or activity. In
 CC particular, they are useful for promoting infant growth due to growth
 CC hormone deficiency. The compounds of the invention are safe with no
 CC accompanying side effects. The present sequence represents a ghrelin-type
 CC growth hormone secretagogue (GHS) precursor protein of the invention
 XX
 SQ Sequence 117 AA;

Query Match 32.1%; Score 198; DB 4; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;
 Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||
 DB 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||

RESULT 10
 AAE23838
 ID AAE23838.standard; protein; 117 AA.
 XX

DB 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||
 DB 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||

RESULT 9
 ABB78319
 ID ABB78319 standard; protein; 117 AA.
 XX
 AC ABB78319;
 XX
 DT 05-DEC-2002 (first entry)
 XX
 DE Amino acid sequence of a human zsig33.
 XX
 KW Short gastrointestinal peptide; SGIP; zsig33; motilin.
 XX
 OS Homo sapiens.
 XX
 FH Key Location/Qualifiers
 FT Peptide 1..23
 FT Protein /note= "signal peptide"
 FT 24..119
 FT /note= "mature protein"
 XX
 PN US6420521-B1.
 XX
 PD 16-JUL-2002.
 XX
 PF 30-JUN-2000; 2000US-00608810.
 XX
 PR 30-JUN-1999; 99US-0141592P.
 XX
 PA (ZYMO) ZYMOGENETICS INC.
 XX
 PI Sheppard PO, Jaspers SR, Deisher TA, Bishop PD;
 XX
 DR WPI: 2002-634794/68.
 DR N-PSDB; ABV72214.
 XX
 PT New Short Gastrointestinal Peptide, which has homology to motilin, useful
 PT for preventing, diagnosing and treating gastrointestinal disorders.
 XX
 PS Disclosure; Col 39-40; 23pp; English.
 XX
 CC The present sequence represents human zsig33. The specification describes
 CC a short gastrointestinal peptide (SGIP), which is derived from zsig33.
 CC SGIP has homology to motilin. The SGIP peptide may be used in the
 CC prevention, diagnosis and treatment of diseases associated with
 CC inappropriate SGIP expression. For example, SGIP may be used to treat
 CC disorders associated with decreased expression by rectifying mutations or
 CC deletions in a patient's genome that affect the activity of SGIP by
 CC expressing inactive proteins or to supplement the patients own production
 CC of SGIP. SGIP may also be used as an antigen in the production of
 CC antibodies against SGIP and in assays to identify modulators of SGIP
 CC expression and activity. The anti-SGIP antibodies, agonists and
 CC antagonists may also be used to regulate expression and activity. The
 CC anti-SGIP antibodies may also be used as diagnostic agents for detecting
 CC the presence of SGIP in samples
 XX
 SQ Sequence 117 AA;

Query Match 32.1%; Score 198; DB 5; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;
 Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||
 DB 1 MPSPGTVCSSLILGLMLDLAMAGSSFLSPHQVQVRPPHKAP 44
 |||||

RESULT 10
 AAE23838
 ID AAE23838 standard; protein; 117 AA.
 XX

AC AAE23838;
 XX DT 10-SEP-2002 (first entry)
 XX DE Human zsig33 protein.
 XX KW Human; zsig33-like peptide; gastric contractility; nutrient uptake;
 KW growth hormone; digestive enzyme; restorative therapy; gene therapy;
 KW protein therapy; gastrointestinal; endocrine; anabolic.
 XX OS Homo sapiens.
 XX PN US2002055156-A1.
 XX PD 09-MAY-2002.
 XX PF 10-MAY-2001; 2001US-00853253.
 XX PR 11-MAY-2000; 2000US-0203300P.
 XX (JASP/) JASPERS S R.
 PA (SHEP/) SHEPPARD P O.
 PA (DEIS/) DEISHER T A.
 PA (BISH/) BISHOP P D.
 XX Jaspers SR, Sheppard PO, Deisher TA, Bishop PD;
 XX WPI; 2002-443750/47.
 XX N-PSDB; AAD38238.
 XX ZSIG33-like peptides and polynucleotides, useful for modulating gastric
 PT contractility, nutrient uptake, growth hormones and/or secretion of
 PT digestive/pancreatic enzymes and hormones.
 XX Disclosure; Page 27; 34pp; English.
 XX The invention relates to zsig33-like peptides and their corresponding
 CC nucleic acids and methods for modulating gastric contractility, nutrient
 CC uptake, growth hormones, secretion of digestive enzymes and hormones. The
 CC sequences of the invention are used in the prevention, diagnosis and
 CC treatment of diseases associated with inappropriate ZSIG33 expression.
 CC The nucleic acids of the invention and their complements are used as DNA
 CC probes in diagnostic assays to detect and quantitate the presence of
 CC similar nucleic acids in samples, and therefore which patients may be in
 CC need of restorative therapy. The ZSIG33 peptides are used as antigens in
 CC the production of antibodies against ZSIG33 and in assays to identify
 CC modulators of ZSIG33 expression and activity. The anti-ZSIG33 antibodies
 CC and antagonists are used to down regulate expression and activity. The
 CC anti-ZSIG33 antibodies are also used as diagnostic agents for detecting
 CC the presence of ZSIG33 in samples (e.g. by enzyme linked immunosorbent
 CC assay (ELISA)). The peptides and nucleic acids of the invention are used
 CC to modulate gastric contractility, nutrient uptake, growth hormones, the
 CC secretion of digestive enzymes and hormones, and/or secretion of enzymes
 CC and/or hormones in the pancreas. zsig33-like DNA is used in gene therapy
 CC and zsig33-like peptide is used in protein therapy. The present sequence
 CC is human zsig33 protein
 XX Sequence 117 AA;
 SQ
 Query Match 32.1%; Score 198; DB 5; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;
 Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;
 QY 1 MPSFGTVCSSLLGLGWLMDLAWAGSSFLSPFHQVQVRPPHKAP 44
 DB 1 MPSFGTVCSSLLGLGWLMDLAWAGSSFLSPFHQVQVRPPHKAP 44
 RESULT 11
 AAE15883
 ID AAE15883 standard; protein; 117 AA.
 XX AAE15883;
 AC AAE15883;
 DT 26-MAR-2002 (first entry)
 DE Human zsig33 protein.
 KW Human; zsig33-like peptide; ZS33LP; immunity; developmental process;
 KW infection; human immunodeficiency virus; vaccine; antihypoglycaemic;
 KW adsorption enhancer; gastrointestinal disease; growth related disease;
 KW inflammation; gene therapy; growth regulation; blood vessel formation;
 KW HIV; zsig33 protein.
 XX OS Homo sapiens.
 XX PH 1. .23
 FT Peptide /label= Signal_peptide
 FT Protein 24. .117
 FT /note= "Human mature zsig33 protein"
 XX WO200187933-A2.
 XX 22-NOV-2001.
 XX 10-MAY-2001; 2001WO-US015091.
 XX 11-MAY-2000; 2000US-00569271.
 XX (ZYMO) ZYMOGENETICS INC.
 XX Jaspers SR, Sheppard PO, Deisher TA, Bishop PD;
 XX WPI; 2002-082982/11.
 XX N-PSDB; AAD25759.
 XX New polypeptides, useful for modulating gastric contractility, nutrient
 PT uptake, pancreatic secretion of hormones, digestive enzymes and treating
 PT gastrointestinal and growth related diseases, comprises zsig33-like
 PT peptides.
 XX Disclosure; Page 80-81; 89pp; English.
 XX The invention relates to zsig33-like peptides (ZS33LP) including zsig33-
 CC linker, zsig33-beta, zsig33-gamma, zsig33-delta and zsig33-epsilon
 CC peptides and nucleic acid molecules encoding such zsig33-like peptides.
 CC ZS33LP peptides activate the immune system in boosting immunity to
 CC infectious diseases, treating immunocompromised patients such as human
 CC immunodeficiency virus (HIV) patients, in improving vaccines and in
 CC treatment of bacterial, viral, protozoal and fungal infections. Peptides
 CC of the invention are used to identify and isolate receptors involved in
 CC growth regulation in the liver, blood vessel formation and other
 CC developmental processes. They are useful for evaluating functions of
 CC hypothalamus-pituitary-adrenal axis, to modulate growth and/or
 CC differentiation of tumour cells, as additives to anti- hypoglycaemic
 CC preparations containing glucose and as adsorption enhancers for oral
 CC drugs which require fast nutrient action and to stimulate glucose-induced
 CC insulin release. They are also useful as research reagents for the
 CC expansion, differentiation, growth factor and hormone secretion and/or
 CC cell-cell interactions of tissues associated with gastrointestinal
 CC system, brain and central nervous system. These molecules are useful for
 CC treating dysfunction associated with contractile tissues or to suppress
 CC or enhance contractility in vivo and to treat gastrointestinal and growth
 CC related diseases. ZS33LP peptides, nucleic acids and/or antibodies are
 CC useful for treating disorders associated with gastrointestinal
 CC contractility, secretion of digestive enzymes, hormone and acids,
 CC secretion of hormones in the pancreas and/or brain, gastrointestinal
 CC motility, recruitment of digestive enzymes, inflammation and regulation
 CC of nutrient absorption. Sequences of the invention are useful in gene
 CC therapy. The present sequence is human zsig33 protein
 XX Sequence 117 AA;
 SQ
 Query Match 32.1%; Score 198; DB 5; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;

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Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;
QY 1 MPEPGTVCSSLLILGLMLWLDLAMAGSSFLSPFHQRVQVRPHKAP 44
    |||||
Db 1 MPEPGTVCSSLLILGLMLWLDLAMAGSSFLSPFHQRVQVRKESKAP 44

RESULT 12
ID ABU58046 standard; protein; 117 AA.
AC ABU58046;
XX
XX 14-APR-2003. (first entry)
XX Human PRO polypeptide #78.
DE
DE Human; PRO; cytostatic; tumour; cancer; breast; lung; stomach; liver;
KW horse; cow; dog; cat; sheep; pig; goat; rabbit; ADEPT;
KW antibody-dependent enzyme mediated prodrug therapy.
XX
XX Homo sapiens.
XX
XX US2003027163-A1.
XX
XX 06-FEB-2003.
XX
XX 15-NOV-2001; 2001US-00997666.
XX
XX 16-JUN-1997; 97US-0049787P.
XX 17-OCT-1997; 97US-0062250P.
XX 05-NOV-1997; 97WO-US020069.
XX 12-NOV-1997; 97US-0065186P.
XX 13-NOV-1997; 97US-0065311P.
XX 24-NOV-1997; 97US-0066770P.
XX 25-FEB-1998; 98US-0075945P.
XX 20-MAR-1998; 98US-0078910P.
XX 28-APR-1998; 98US-0083322P.
XX 07-MAY-1998; 98US-0084500P.
XX 28-MAY-1998; 98US-0087106P.
XX 02-JUN-1998; 98US-0087607P.
XX 02-JUN-1998; 98US-0087609P.
XX 02-JUN-1998; 98US-0087759P.
XX 03-JUN-1998; 98US-0087827P.
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XX 04-JUN-1998; 98US-0088025P.
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XX 04-JUN-1998; 98US-0088029P.
XX 04-JUN-1998; 98US-0088030P.
XX 04-JUN-1998; 98US-0088033P.
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XX 10-JUN-1998; 98US-0088734P.
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XX 11-JUN-1998; 98US-0088861P.
XX 11-JUN-1998; 98US-0088876P.
XX 12-JUN-1998; 98US-0089105P.
XX 16-JUN-1998; 98US-0089440P.
XX 16-JUN-1998; 98US-0089512P.
XX 16-JUN-1998; 98US-0089514P.
XX 17-JUN-1998; 98US-0089532P.
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XX 17-JUN-1998; 98US-0089598P.
XX 17-JUN-1998; 98US-0089599P.
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XX 17-JUN-1998; 98US-0089601P.
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XX 18-JUN-1998; 98US-0089608P.
XX 19-JUN-1998; 98US-0089947P.
XX 19-JUN-1998; 98US-0089948P.
XX 19-JUN-1998; 98US-0089952P.
XX 22-JUN-1998; 98US-0090246P.
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XX 23-JUN-1998; 98US-0090349P.
XX 23-JUN-1998; 98US-0090355P.
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XX 24-JUN-1998; 98US-0090444P.
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XX 25-JUN-1998; 98US-0090694P.
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XX 26-JUN-1998; 98US-0090696P.
XX 26-JUN-1998; 98US-0090862P.
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XX 02-JUL-1998; 98US-0091544P.
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XX 02-JUL-1998; 98US-0091626P.
XX 02-JUL-1998; 98US-0091628P.
XX 02-JUL-1998; 98US-0091633P.
XX 02-JUL-1998; 98US-0091646P.
XX 02-JUL-1998; 98US-0091673P.
XX 07-JUL-1998; 98US-0091978P.
XX 07-JUL-1998; 98US-0091982P.
XX 09-JUL-1998; 98US-0092182P.
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XX 30-JUL-1998; 98US-0093339P.
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XX 11-AUG-1998; 98US-0096143P.
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XX 12-AUG-1998; 98US-0096329P.
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XX 17-AUG-1998; 98US-0096895P.
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XX 18-AUG-1998; 98US-0096950P.
XX 18-AUG-1998; 98US-0096959P.
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PR	18-AUG-1998;	98US-0096960P.	RESULT 13	
PR	18-AUG-1998;	98US-0097022P.	ABUS9124	
PR	19-AUG-1998;	98US-0097141P.	ID ABUS9124 standard; protein; 117 AA.	
PR	20-AUG-1998;	98US-0097218P.	XX	
PR	20-AUG-1998;	98US-0097661P.	AC ABUS9124;	
PR	26-AUG-1998;	98US-0097952P.	XX	
PR	26-AUG-1998;	98US-0097954P.	DT 28-APR-2003 (first entry)	
PR	26-AUG-1998;	98US-0097955P.	XX	
PR	26-AUG-1998;	98US-0097974P.	DE	Novel human secreted or transmembrane protein PRO1066.
PR	26-AUG-1998;	98US-0097978P.	XX	
PR	26-AUG-1998;	98US-0097979P.	KW	Human; PRO; hypertrophy of neonatal heart; angiogenesis; wound healing;
PR	26-AUG-1998;	98US-0097986P.	KW	cardiac insufficiency disorder; cancer; tumor; immune response;
PR	31-AUG-1998;	98US-0098014P.	KW	adrenal cortical capillary endothelial growth; c-fos induction;
PR	16-SEP-1998;	98US-0098525P.	KW	vascular endothelial growth factor inhibition; VEGF inhibition;
PR	16-SEP-1998;	98US-0100634P.	KW	endothelial cell growth inhibitor; T-lymphocytes stimulation;
PR	17-SEP-1998;	98US-0100634P.	KW	retinal neurons cell survival; rod photoreceptor cell survival;
PR	17-SEP-1998;	98US-0100634P.	KW	retinal disorder; retinitis pigmentosa; kidney disorder;
PR	17-SEP-1998;	98US-0100634P.	KW	mammalian kidney mesangial cell proliferation; Berger disease;
PR	17-SEP-1998;	98US-0100634P.	KW	dermatitis; herpeticiformis; Crohn's disease; chondrocyte proliferation;
PR	07-OCT-1998;	98US-0100634P.	XX	chondrocyte redifferentiation; sports injury; arthritis.
PR	01-DEC-1998;	98US-0100634P.	OS	Homo sapiens.
PR	01-DEC-1998;	98US-0100634P.	XX	
PR	01-DEC-1998;	98US-0100634P.	PN	US2002132252-A1.
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PR	01-DEC-1998;	98US-0100634P.	PD	19-SEP-2002.
PR	01-DEC-1998;	98US-0100634P.	XX	
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PR	01-DEC-1998;	98US-0100634P.	XX	
PR	01-DEC-1998;	98US-0100634P.	PR	16-JUN-1997; 97US-0049787P.
PR	01-DEC-1998;	98US-0100634P.	PR	17-OCT-1997; 97US-0062250P.
PR	01-DEC-1998;	98US-0100634P.	PR	05-NOV-1997; 97US-0062250P.
PR	01-DEC-1998;	98US-0100634P.	PR	12-NOV-1997; 97US-0065186P.
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PR	01-DEC-1998;	98US-0100634P.	PR	25-FEB-1998; 98US-0075945P.
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PR	01-DEC-1998;	98US-0100634P.	PR	16-JUN-1998; 98US-0089440P.
PR	01-DEC-1998;	98US-0100634P.	PR	16-JUN-1998; 98US-0089512P.
PR	01-DEC-1998;	98US-0100634P.	PR	16-JUN-1998; 98US-0089514P.
PR	01-DEC-1998;	98US-0100634P.	PR	17-JUN-1998; 98US-0089532P.
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Query Match 32.1%; Score 198; DB 6; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;
 Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;
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 Db 1 MPSGTGCSLLLLGMLWLDLAMAGSSFLSPHQVQVRPPHKAP 44

PR 17-JUN-1998; 98US-0089599P.
 PR 17-JUN-1998; 98US-0089600P.
 PR 17-JUN-1998; 98US-0089653P.
 PR 18-JUN-1998; 98US-0089801P.
 PR 18-JUN-1998; 98US-0089907P.
 PR 18-JUN-1998; 98US-0089908P.
 PR 18-SEP-1998; 98US-0019330.
 PR 17-SEP-1998; 98WO-US019437.
 PR 07-OCT-1998; 98WO-US021141.
 PR 01-DEC-1998; 98WO-US025108.
 PR 05-JAN-1999; 99WO-US000108.
 PR 08-MAR-1999; 99WO-US005028.
 PR 02-JUN-1999; 99WO-US012252.
 PR 15-SEP-1999; 99WO-US021090.
 PR 30-NOV-1999; 99WO-US021547.
 PR 01-DEC-1999; 99WO-US028313.
 PR 01-DEC-1999; 99WO-US028301.
 PR 16-DEC-1999; 99WO-US028634.
 PR 20-DEC-1999; 99WO-US030095.
 PR 06-JAN-2000; 99WO-US030911.
 PR 06-JAN-2000; 2000WO-US000376.
 PR 11-FEB-2000; 2000WO-US003565.
 PR 18-FEB-2000; 2000WO-US004341.
 PR 22-FEB-2000; 2000WO-US004414.
 PR 24-FEB-2000; 2000WO-US004914.
 PR 24-FEB-2000; 2000WO-US005004.
 PR 02-MAR-2000; 2000WO-US005841.
 PR 10-MAR-2000; 2000WO-US006319.
 PR 15-MAR-2000; 2000WO-US006884.
 PR 20-MAR-2000; 2000WO-US007377.
 PR 30-MAR-2000; 2000WO-US008439.
 PR 15-MAY-2000; 2000WO-US013358.
 PR 17-MAY-2000; 2000WO-US013705.
 PR 22-MAY-2000; 2000WO-US014042.
 PR 30-MAY-2000; 2000WO-US014941.
 PR 02-JUN-2000; 2000WO-US015264.
 PR 28-JUL-2000; 2000WO-US020710.
 PR 11-AUG-2000; 2000WO-US022031.
 PR 23-AUG-2000; 2000WO-US023522.
 PR 24-AUG-2000; 2000WO-US023328.
 PR 08-NOV-2000; 2000WO-US030952.
 PR 01-DEC-2000; 2000WO-US032678.
 PR 28-FEB-2001; 2001WO-US006520.
 PR 01-JUN-2001; 2001WO-US017800.
 PR 20-JUN-2001; 2001WO-US019692.
 PR 28-JUN-2001; 2001WO-US021066.
 PR 09-JUL-2001; 2001WO-US021735.
 PR 28-AUG-2001; 2001US-00941992.
 (GETH) GENENTECH INC.
 PI Ashkenazi AJ, Baker KP, Botstein D, Desnoyers L, Eaton DL;
 PI Ferrara N, Fong S, Gerber H, Gerritsen ME, Goddard A, Godowski PJ;
 PI Grimaldi JC, Gurney AL, Kijavini IJ, Napier MA, Pan J, Paoni NF;
 PI Roy MA, Stewart TA, Tumas D, Watanabe CK, Williams PM, Wood WI,
 PI Zhang Z;
 XX WPI; 2003-247083/24.
 DR N-PSDB; ABX80294.
 XX
 PT Novel isolated PRO polypeptides e.g., PRO826, PRO1068, PRO1184, PRO1346
 PT and PRO1375, which stimulate proliferation of stimulated T-lymphocytes
 PT are therapeutically useful for enhancing immune response and in cancer
 PT treatments.
 XX
 PS Claim 12; Fig 186; 649pp; English.
 XX
 CC The invention describes an isolated human PRO polypeptide. The PRO
 CC polypeptides are useful in detecting PRO polypeptides in a sample, in
 CC linking a bioactive molecule to a cell expressing a PRO polypeptide, and
 CC in modulating at least one biological activity of a cell expressing a PRO
 CC polypeptide. PRO1312 stimulates hypertrophy of neonatal heart and is thus

CC useful for treating cardiac insufficiency disorders. PRO1154 and PRO1186
 CC stimulate adrenal cortical capillary endothelial growth, and PRO336,
 CC PRO943, PRO828, PRO826, PRO1068 or PRO535, PRO826, PRO819, PRO1126,
 CC PRO1360 and PRO1387 induce c-fos in endothelial cells, and are thus
 CC useful for treating conditions or disorders where angiogenesis would be
 CC beneficial, e.g. wound healing and antagonist of this polypeptide are
 CC useful for treating cancerous tumours. PRO812 inhibits vascular
 CC endothelial growth factor (VEGF) stimulated proliferation of endothelial
 CC cells and is thus useful for inhibiting endothelial cell growth in
 CC mammals which would be beneficial in inhibiting tumour growth. PRO826,
 CC PRO1068, PRO1184, PRO1346 and PRO1375 stimulate proliferation of
 CC stimulated T-lymphocytes and are therapeutically useful for enhancing
 CC immune response. PRO828, PRO826, PRO1068 or PRO1132 enhance survival of
 CC retinal neurons cells (PRO1132 is also enhances survival/proliferation of
 CC rod photoreceptor cells) and therefore are useful for treating retinal
 CC disorders of injuries, e.g. retinitis pigmentosa, AMD. PRO819, PRO813
 CC and PRO1066 induce proliferation of mammalian kidney mesangial cells,
 CC and therefore are useful for treating kidney disorders associated with
 CC decreased mesangial cell function such as Berger disease or other
 CC nephropathies associated with dermatitis, herpeticiformis or Crohn's
 CC disease. PRO1310, PRO844, PRO1312, PRO1192 and PRO1387 induce the
 CC proliferation and/or redifferentiation of chondrocytes in culture and are
 CC thus useful for treating sports injuries, and arthritis. This is the
 CC amino acid sequence of a novel human PRO protein
 XX
 SQ Sequence 117 AA;
 Query Match 32.1%; Score 198; DB 6; Length 117;
 Best Local Similarity 88.6%; Pred. No. 3.9e-14;
 Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;
 QY 1 MPSPGTVCVCSLLLLGLMLDLAMAGSSFLSPHQVQVRPPHAP 44
 DB 1 MPSPGTVCVCSLLLLGLMLDLAMAGSSFLSPHQVQVRPPHAP 44
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 ABU82636
 ID ABU82636 standard; protein; 117 AA.
 XX AC ABU82636;
 XX DT 26-JUN-2003 (first entry)
 XX DE Human secreted/transmembrane protein PRO1066.
 XX KW Human; PRO; secreted protein; transmembrane protein;
 KW cardiac insufficiency disorders; angiogenesis; wound healing;
 KW cancerous tumour; immune response; retinal disorder; sight loss;
 KW retinitis pigmentosa; age-related macular degeneration; AMD;
 KW kidney disorder; Berger disease; nephropathy; dermatitis; herpeticiformis;
 KW Crohn's disease; sports injury; arthritis.
 XX OS Homo sapiens.
 XX PN US2003032023-A1.
 XX PD 13-FEB-2003.
 XX PF 14-NOV-2001; 2001US-00990711.
 XX PR 16-JUN-1997; 97US-0049787P.
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Query Match 32.1%; Score 198; DB 6; Length 117;
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RESULT 15
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ID ABO17836 standard; protein; 117 AA.
XX
AC ABO17836;
XX
DT 26-AUG-2003 (first entry)
XX
DE Novel human secreted and transmembrane protein PRO1066.
XX
KW Human; secreted and transmembrane protein; PRO; antiinflammatory;
KW antiarteriosclerotic; cardiant; anti-infertility; anti-HIV; cytostatic;
KW antidiabetic; gene therapy; tumour necrosis factor (TNF)-alpha release;
KW TNF-alpha release; cell proliferation; cell differentiation;
KW gene expression modulator; proteoglycan release; cytokine release;
KW tumour; inflammatory disease; organ failure; atherosclerosis;
KW cardiac injury; infertility; birth defect; premature aging; AIDS;
KW acquired immunodeficiency syndrome; cancer; diabetic complication;
KW chromosome mapping; gene mapping; pharmaceutical; diagnostic; biosensor;
KW bioreactor; tissue typing.
XX
OS Homo sapiens.
XX
PN US2003032156-A1.
XX
PD 13-FEB-2003.
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PF 06-MAY-2002; 2002US-00140474.
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PR 30-NOV-1999; 99WO-US028409.
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XX (GETH) GENENTECH INC.
XX PA
XX PI Baker KP, Beresini M, Deforge L, Desnoyers L, Filvaroff E, Gao W;
XX PI Gerritsen ME, Goddard A, Godowski FJ, Gurney AL, Sherwood S;
XX PI Smith V, Stewart TA, Tumas D, Watanabe CK, Wood WI, Zhang Z;
XX DR N-PSDB; ACD24073.
XX PS WPI; 2003-341980/32.
XX DR N-PSDB; ACD24073.
XX PT New secreted and transmembrane PRO nucleic acids, for treating
XX PT inflammation, organ failure, atherosclerosis, cardiac injury,
XX PT infertility, birth defects, premature aging, acquired immunodeficiency
XX PT syndrome (AIDS), or cancer.
XX PS
XX PS Claim 12; Fig 442; 660pp; English.
XX CC The invention describes an isolated nucleic acid (I) comprising, or which
XX CC has 80 % sequence identity to, or the full-length coding sequence of, one
XX CC of 275 nucleotide sequences, and which encodes a corresponding
XX CC polypeptide selected from 275 amino acid sequences, where all sequences
XX CC are given in the specification. The polypeptide encoded by (I) is used to
XX CC detect PRO polypeptides, link a bioactive molecule to a cell expressing a
XX CC PRO polypeptide, modulate a biological activity of a cell, stimulate the
XX CC release of tumour necrosis factor (TNF)-alpha from human blood, modulate
XX CC the uptake of glucose or free fatty acid by cells, stimulate or inhibit
XX CC the proliferation or differentiation of cells or gene expression,
XX CC stimulate the release of proteoglycans, stimulate the release of cytokine
XX CC from peripheral blood mononuclear cells, inhibit the binding of a peptide
XX CC to factor VIIA, or detect the presence of tumour in a mammal. The nucleic
XX CC acid and polypeptide encoded by it, are useful for treating inflammatory
XX CC diseases, organ failure, atherosclerosis, cardiac injury, infertility,
XX CC birth defects, premature aging, acquired immunodeficiency syndrome
XX CC (AIDS), cancer, or diabetic complications. The nucleic acid is useful as
XX CC hybridisation probes, in chromosome and gene mapping, and in generating
XX CC antisense RNA or DNA. The polypeptides are useful as pharmaceuticals,
XX CC diagnostics, biosensors or bioreactors. Both are useful in tissue typing.
XX CC This is the amino acid sequence of a novel human secreted and
XX CC transmembrane PRO polypeptide
XX SQ Sequence 117 AA;
Query Match 32.1%; Score 198; DB 6; Length 117;
Best Local Similarity 88.6%; Pred. No. 3.9e-14;
Matches 39; Conservative 0; Mismatches 5; Indels 0; Gaps 0;
OY 1 MPSPGTVCISLLGLMLDLAMAGSSFLSPHQHVQVRPPKAP 44
DB 1 MPSPGTVCISLLGLMLDLAMAGSSFLSPHQHVQVRPPKAP 44

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